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Are autosomal sex-determining factors of the housefly (*Musca domestica*) spreading north?

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Summary

Multiple sex-determining factors have been found in natural populations of the housefly, *Musca domestica*. Their distribution seems to follow a geographical cline. The ‘standard’ system, with a male-determining factor, *M*, located on the Y chromosome, prevails at higher latitudes and altitudes. At lower latitudes and altitudes *M* factors have also been found on any of the five autosomes. Such populations often also harbour a dominant autosomal factor, *F^D*, which induces female development even in the presence of several *M* factors. Autosomal *M* factors were first observed some 50 years ago. It has been hypothesized that following their initial appearance, they are spreading northwards, replacing the standard XY system, but this has never been systematically investigated. To scrutinize this hypothesis, we here compare the current distribution of autosomal *M* factors in continental Europe, on a transect running from Germany to southern Italy, with the distribution reported 25 years ago. Additionally, we analysed the frequencies of the *F^D* factor, which has not been done before for European populations. In contrast to earlier predictions, we do not find a clear change in the distribution of sex-determining factors: as 25 years ago, only the standard XY system is present in the north, while autosomal *M* factors and the *F^D* factor are prevalent in Italy. We discuss possible causes for this apparently stable polymorphism.

1. Introduction

Sex determination in the housefly, *Musca domestica*, is more variable than in most other species, which usually exhibit just a single sex-determining mechanism (Bull, 1983; Dübendorfer *et al.*, 2002). Polymorphism for sex-determining factors has been found in many natural populations of the housefly (Franco *et al.*, 1982; Denholm *et al.*, 1985; Tomita & Wada, 1989b; Feldmeyer *et al.*, submitted; Table 1). In ‘standard’ strains, sex is determined by a male-determining factor, *M*, which is located on the Y chromosome; therefore males are XY and females are XX. During development, the *M* factor blocks the female-determining factor *F* located on autosome IV, the activity of which is necessary for female development. In many populations, *M* is located on one of the autosomes or even on the X chromosome (Denholm

et al., 1983). In such populations, usually a dominant constitutive mutation of *F* (*F^D*) is also present, which triggers female development even in the presence of several *M* factors in the same individual (see McDonald *et al.*, 1978; Franco *et al.*, 1982; Dübendorfer *et al.*, 2002; Table 1).

The XY system is probably ancestral in the housefly, since it is also very common in closely related species (Boyes *et al.*, 1964) and the first reports on autosomal sex-determining (SD) factors appeared only around 1960 (reviewed by Franco *et al.*, 1982). Since then, the geographical distribution of different SD factors has been studied on most of the continents and appears to follow geographical clines. In general, the Y chromosome is more common at higher latitudes and altitudes and its frequency gradually decreases with decreasing latitude and altitude, leading to populations with only autosomal sex-determining factors (autosomal *M* and *F^D*) closer to the equator

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Table 1. Relation between genotype and gender in the housefly

Autosomes		Sex chromosomes	
IV	I–V	XX	XY
F/F	$+/+$	♀	♂
F/F	$M/*$	♂	♂
F/F^D	$*/*$	♀	♀

The female-determining factors (F/F^D) are located on autosome IV. The male-determining factors (M) can be located on any chromosome. A '+' indicates the wild-type state (no M) and a '*' indicates that an M or + allele on this locus will not influence the sex.

and at low altitudes (Franco *et al.*, 1982; Tomita & Wada, 1989b; Çakir & Kence, 1996; Hamm *et al.*, 2005; Feldmeyer *et al.*, submitted). It is not clear what forces are responsible for the distribution of different SD factors, but temperature seems to be an important factor (Feldmeyer *et al.*, submitted).

There is some evidence that autosomal sex-determining factors have spread in some populations replacing the standard XY system (Franco *et al.*, 1982; Tomita & Wada, 1989a,b). It has been hypothesized (Franco *et al.*, 1982; Denholm *et al.*, 1985; Tomita & Wada, 1989a,b; Çakir & Kence, 1996) that the observed distributions are a transient state. In particular, Franco and colleagues (1982) suggested that autosomal M factors are spreading north in Europe, but their hypothesis was based only on the change in frequency of the Y chromosome in a few populations before 1980. No systematic or recent studies have been done on the dynamics of different SD factors in natural populations of the housefly. The last study in continental Europe dates from 25 years ago (Franco *et al.*, 1982), in which cytological data were used to show a clear latitudinal cline with the standard XY system exclusively present in the north of Europe (Iceland, Denmark, the Netherlands, Germany and Switzerland) and entirely autosomal populations (lacking the Y chromosome) in southern Italy at altitudes below 100 m. In northern Italy mixed populations have been found, with the frequency of the Y chromosome increasing with higher altitudes and latitudes.

The aim of this study was to investigate whether the distribution of SD factors in the housefly has changed in Europe over the last 25 years. Therefore, we sampled a number of European populations on a north–south transect from Germany to southern Italy, and compared the frequency of males that carry the Y chromosome and autosomal M factors with the data published by Franco and colleagues (1982). Additionally, we analysed the frequencies of the F^D factor and we publish the frequencies of M factors

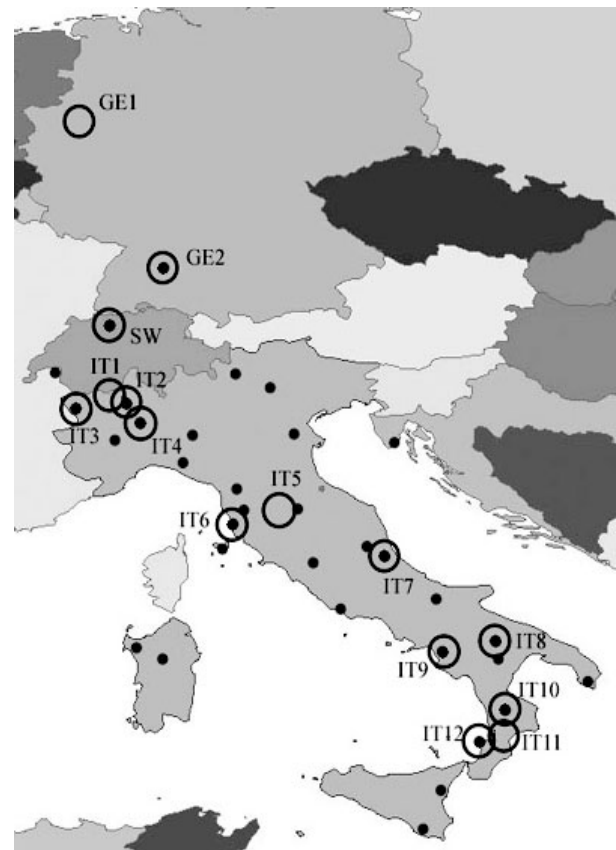


Fig. 1. Sampling locations in the study of Franco and colleagues (1982; dots) and in the present study (circles). Locations from the present study are labelled with population codes as in Table 2.

located on different chromosomes, which has not been done before for European housefly populations.

2. Materials and methods

(i) Collection and rearing of flies

We sampled populations along a north–south transect from north Germany to south Italy in July 2006 (see Fig. 1 and Table 2 for details on the sampling locations). Most of the sampling sites were chosen to be close to the ones studied by Franco *et al.* (1982), as far as we could judge from the limited information available. For Germany and Switzerland, they gave only the name of a state (Baden-Württemberg) or a canton (Mittelland) and our sampling sites lie within these areas. For Italy, Franco and colleagues published a map indicating sampling sites together with information on altitudes, but precise geographical coordinates were lacking. We judged their locations visually and used altitudes within 110 m, but usually within a 50 m range. The exception is population IT5 where the altitude given by Franco *et al.* (1982) does not match the area indicated by them, so to match the altitude we sampled 50 km west of their indicated location. Ultimately, our sampling sites were

Table 2. *Geographical coordinates, altitudes (in metres above sea level) and average yearly temperatures of the sampling sites*

Population code	Latitude (°N)	Longitude (°E)	Altitude (m)	Temperature (°C)
GE1	51° 19·4'	7° 10·9'	220	9·1
GE2	48° 29·5'	9° 2·0'	347	9·0
SW	47° 17·8'	7° 51·8'	410	9·4
IT1	45° 46·6'	8° 2·5'	794	8·8
IT2	45° 42·3'	8° 14·1'	470	10·1
IT3	45° 35·4'	7° 8·0'	1700	4·2
IT4	45° 17·8'	8° 33·1'	121	12·3
IT5	43° 29·2'	11° 33·1'	313	13·2
IT6	43° 11·0'	10° 31·7'	18	15·4
IT7	42° 32·6'	13° 49·3'	367	13·3
IT8	40° 45·7'	16° 14·3'	562	13·3
IT9	40° 32·5'	15° 6·4'	63	16·1
IT10	39° 21·4'	16° 26·5'	1194	10·4
IT11	38° 48·0'	16° 20·3'	690	13·9
IT12	38° 40·6'	15° 54·6'	49	17·7

Sampling sites are ordered according to their latitude. Letters in the code indicate the country of origin: GE, Germany; SW, Switzerland; IT, Italy.

distributed approximately homogeneously along a north–south transect, with some areas having sampling sites at different altitudes.

For each location, we obtained data on average monthly minimum and maximum temperatures from WORLDCLIM (www.worldclim.org; see Hijmans *et al.*, 2005), which provides global estimates at a resolution of 1 km². We estimated average yearly temperatures as the mid-point between minimum and maximum temperatures (Table 2). Since all these measures of temperature are highly correlated ($P < 0.0001$, Pearson's product-moment correlation test), we used only the average yearly temperature in our statistical analysis (see below).

Flies were sampled at farms and horse stables. At each location we caught approximately 50 adult males and females (except for IT3, where only 10 females were found). The flies were caught with sweeping nets, placed in plastic containers and provided with water and milk powder as food. They were also provided with egg-laying medium (according to Hilfiker-Kleiner *et al.*, 1994) on which females laid eggs within a few days. Larvae were transferred to larger containers after a few days and fed *ad libitum* on the same medium. Flies from all the locations (or their offspring) were successfully transported to the laboratory and populations were established and maintained in cages at a population size of approximately 500 individuals.

(ii) Analysis of the sex-determining factors

(a) *M factors*. The presence of different *M* factors in males was determined by two generations of

single-pair crosses with standard XX (without an F^D factor) virgin females, from a marker strain that carries visible recessive mutations in the homozygous state on each of the five autosomes (Tomita & Wada, 1989*b*). The sex ratio of F1 offspring shows whether the father was homozygous for at least one *M* factor (only sons are produced) or heterozygous for all *M* factors (daughters are also present among the offspring). Sex-linked inheritance of visible markers in the second generation of backcrosses to marker-strain females shows on which chromosomes *M* factors are located. This is a standard procedure in our laboratory and it gives a good estimation of the frequency of *M* factors located on different autosomes (for details see Denholm *et al.*, 1983). However, if a focal wild-type male was homozygous for *M* (producing all-male offspring) and all his sons appeared to have two (or more) *M* factors (e.g. *M* on autosome II and V), we could not unambiguously determine whether the father was homozygous for *M* on only one or on both chromosomes, especially if the number of sons was small. For example, M^{II}/M^{II} ; M^V/M^V , $M^{II}/+$; M^V/M^V and M^{II}/M^{II} ; $M^V/+$ males all produce $M^{II}/+$; $M^V/+$ sons when mated with standard females. This happened a few times (13 males in total, with a maximum of 4 males per population). For each chromosome involved in a population, we calculated both the minimal frequency of *M* (assuming that all ambiguous males were heterozygous for *M*) and the maximal frequency of the *M* factor (assuming that all ambiguous males were homozygous for *M*) on the given chromosome. We then used the mid-point value between the two extremes as a population estimate.

We used 20 males from each population for the first series of crosses and 3 sons from each of them for the F1 backcrosses (although we did not obtain offspring from all males). Males used for analysis were either the ones caught in the field (IT3), or from the first generation in the laboratory (offspring of the wild-caught flies: IT6, IT7, IT8, IT10, IT11, IT12), the third generation in the laboratory (GE1, GE2, SW, IT1, IT2, IT4, IT5) or the fourth generation (IT9). Because of the lack of visible markers on the X and the Y chromosomes, in cases in which we assigned *M* to a sex chromosome, we cannot be sure whether it was located on the Y or the X (as has been found in Britain: Denholm *et al.*, 1983, 1985). If *M* was located on a sex chromosome we will call this chromosome Y, but we will discuss this issue in more detail later.

(b) F^D factor. *F* and F^D factors have been sequenced at the University of Zürich (M. Hediger and D. Bopp, personal communication). F^D has two deletions compared with *F* in all populations analysed (of European, Asian and African origin). We used

primers designed for one of these deletions to distinguish between F (one band present) and F^D (two bands) females. We used approximately 20 females from each population, either females caught in the field (populations GE2, SW, IT5 and all 10 females from IT3) or from the first generation in the laboratory (all the other populations). Additionally, we took 2 or 3 females from each population and crossed them individually with a male homozygous for M located on autosome III. Females without F^D produce only sons, but those with F^D also produce daughters, because F^D is dominant over M . After determining the sex of the offspring, we also analysed the mothers molecularly and found without exception that the results of the molecular analysis were consistent with those obtained from the crosses. This shows that the deletion in the F^D factor is also present in the populations we collected and justifies the use of the molecular technique for analysing frequencies of F^D in our populations.

(iii) Statistical analysis

We performed a logistic regression analysis using the glm function with quasi-binomial errors in R (R Development Core Team, 2006) to investigate the influence of latitude, altitude and temperature on the frequency of autosomal M males (with at least one autosomal M factor) and on the frequency of females with the F^D factor. We started with a full model (including all two-way interactions between explanatory variables) and used backward selection to find the minimal adequate model. The significance of the difference between models was assessed with the likelihood-ratio approach, using F -tests to correct for under- and overdispersion (Krackow & Tkadlec, 2001).

A statistical comparison between the frequencies of different SD factors in the past and present is only possible to a limited extent, since Franco *et al.* (1982) performed only cytological observations. They used the frequency of XX males as a measure for the frequency of autosomal males. They checked the linkage of autosomal M factors with crosses similar to ours, but they do not provide the exact frequencies of different factors. They also do not provide data on frequencies of the F^D factor. Moreover, due to the lack of data on the number of males tested by Franco and colleagues (1982), in each autosomal and standard population separately (except for GE2), we could only include eight populations (GE2, IT2, IT3, IT4, IT5, IT7, IT8 and IT10) in a statistical analysis to compare frequencies of autosomal males (without a Y chromosome) between our study and theirs. For this analysis, we performed a mixed-model logistic regression analysis in R using the lmer function with binomial errors from the lme4 package. The full

model included population as a random effect and 'study' (Franco *et al.*, 1982 or this study) as a fixed effect. Significance of the effect of 'study' was judged using the likelihood-ratio approach, using an F -test to correct for overdispersion (Krackow & Tkadlec, 2001). For each of the eight populations we also performed a binomial test, to see whether there is a significant change in the frequency of XX males between the past and the present.

3. Results

(i) Distribution of sex-determining factors in 2006

We found M factors on the sex chromosomes and on each of the autosomes (Table 3, Fig. 2a). M located on autosome III was the most frequent among autosomal M factors and the frequencies of M on autosomes IV and V were very low. We did not detect any autosomal M in the German and Swiss populations or in one northern Italian population from the highest altitude (IT3). In populations with autosomal SD factors, often single males with multiple M factors, located on up to four different chromosomes, were observed (data not shown). Statistical analysis showed that altitude, latitude, temperature and interaction of temperature and latitude (and to a lesser extent interaction between temperature and altitude) influence the frequencies of autosomal M males (Table 4).

We did not find F^D in populations in Germany and only at low frequencies in Switzerland and at the highest location in northern Italy (IT3; Table 3, Fig. 2b). In most of the Italian populations frequencies of F^D females were above 0.75, and in three populations F^D appeared to be at fixation. Statistical analysis showed that the frequency of females with F^D is influenced by latitude, temperature and the interaction of the two (Table 4).

(ii) Comparison with the past

A comparison between our results and those of Franco and colleagues (1982) shows that there is no clear evidence for the spread of autosomal M factors northwards during the last 25 years (Fig. 3). In the two northernmost populations and in IT3, which lacked XX males in the past, we also did not find any autosomal M factor. Furthermore, all populations described by Franco and colleagues (1982) as mixed or autosomal were found to have autosomal M factors in 2006. However, in the populations which were described by Franco and colleagues as autosomal in 1982 (IT6, IT9 and IT12) we also found M on a sex chromosome. Statistical analysis based on the eight populations for which comparable data were available shows no significant systematic change in the

Table 3. Estimated frequencies of females with F^D factor and frequencies of M factors in males in samples from different housefly populations

Population code	No. of females	Frequency of females with F^D	No. of males	Frequency of M on:					
				sex chromosome	autosome I	autosome II	autosome III	autosome IV	autosome V
GE1	20	0.00	18	0.50	0	0	0	0	0
GE2	19	0.00	20	0.50	0	0	0	0	0
SW	21	0.05	20	0.50	0	0	0	0	0
IT1	20	0.44	20	0.52	0	0	0.12	0	0
IT2	21	0.43	16	0.44	0	0.25	0.09	0	0
IT3	10	0.10	11	0.50	0	0	0	0	0
IT4	20	1.00	19	0.42	0.12	0.09	0.45	0	0
IT5	22	1.00	20	0.62	0.02	0.17	0.50	0	0.09
IT6	20	0.95	19	0.68	0.03	0.13	0.32	0	0
IT7	23	0.78	18	0.17	0	0.03	0.53	0	0
IT8	22	1.00	19	0.16	0.09	0.03	0.86	0.03	0.03
IT9	22	0.86	18	0.06	0.08	0.17	0.46	0	0
IT10	19	0.95	19	0	0.03	0	0.55	0.03	0
IT11	23	0.96	17	0.03	0	0	0.76	0	0
IT12	19	0.47	18	0.08	0	0	0.56	0	0

Frequencies of M are given separately for each chromosome (a value of 1.0 would indicate complete homozygosity for M on this chromosome). The sum of M frequencies over all chromosomes may exceed 1.0 when males carry multiple M factors. Population codes are as in Table 2 and Fig. 1.

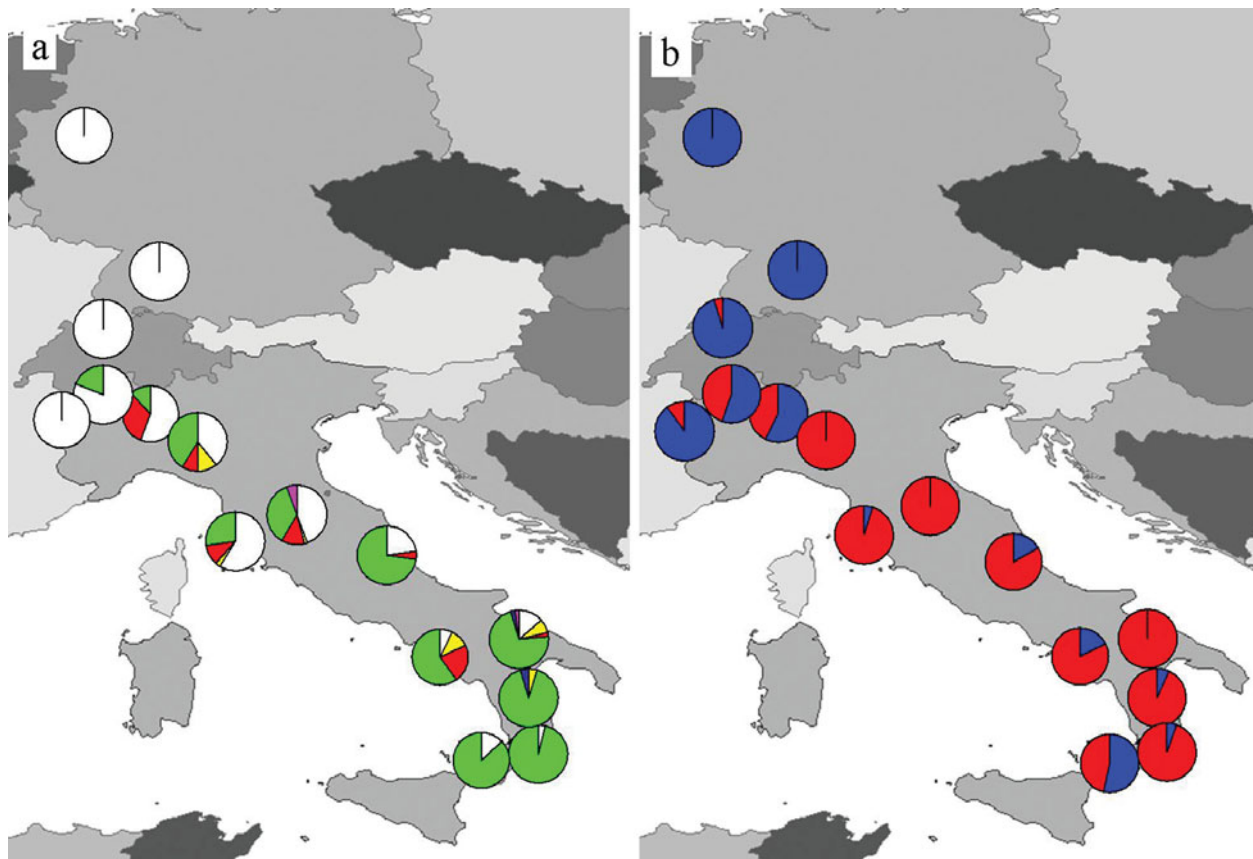


Fig. 2. Distribution of sex-determining factors in the housefly in 2006. (a) Relative frequencies of M factors located on different chromosomes: white, sex chromosome; yellow, autosome I; red, autosome II; green, autosome III; blue, autosome IV; pink, autosome V. (b) Frequencies of females with (red) and without (blue) the F^D factor.

Table 4. Logistic regression analysis of (a) frequencies of autosomal M males and (b) frequencies of females with F^D

Source of variation	Parameter	SE	Δ dev	F	P
(a) Males					
Intercept	277.4	33.2			
Altitude (A)	-0.014	0.002	19.12	70.6	<0.0001
Latitude (L)	-5.521	0.652	28.39	104.9	<0.0001
Temperature (T)	-12.07	1.493	24.24	89.6	<0.0001
A*T	0.0004	0.0002	1.53	5.6	0.042
L*T	0.222	0.029	22.28	82.3	<0.0001
(b) Females					
Intercept	124.470	27.495			
Latitude	-2.884	0.606	108.25	40.95	<0.0001
Temperature	-8.684	1.822	82.04	31.04	<0.0005
L*T	0.204	0.042	85.32	32.28	<0.0005

Parameter estimates (logit scale) and their standard errors (SE) are shown for the final models, after the removal of non-significant variables.

Temperature refers to the average yearly temperature.

Δ dev indicates the change in deviance resulting from removing the given variable from the final model. The F -tests for significance of removed variables have 1 and residual degrees of freedom of the final model (DF) for numerator and denominator, respectively.

Final models: (a) deviance=3.05, residual DF=9; (b) deviance=27.28, residual DF=11.

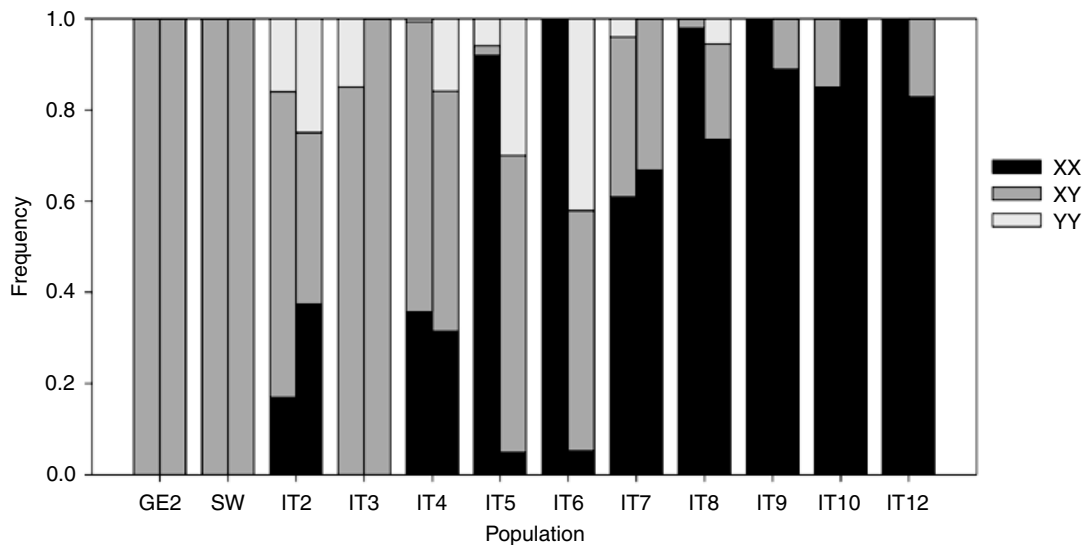


Fig. 3. Comparison of karyotype frequencies in males in the past and the present (2006). For each population the left-hand bar corresponds to the data from Franco *et al.* (1982) and the right-hand bar to the data from this study. We inferred karyotypes from our crosses assuming that Y is the sex chromosome bearing the M factor (see Section 2). Three populations analysed by us are not included in the figure since they were not studied by Franco and colleagues. Populations are ordered according to decreasing latitude of the sampling sites (see Table 2).

frequencies of autosomal males in recent decades (Table 5). Statistical analysis for each population separately, shows a significant decrease in the frequency of XX males for two populations: IT5 and IT8 ($P < 0.002$, which is also significant after Bonferroni correction for multiple tests).

The distribution of F^D also seems to be relatively stable over time. F^D frequencies were not analysed by Franco *et al.* (1982), but the presence of F^D can be

deduced from the occurrence of at least one homozygous M male in all autosomal populations and the occurrence of XY females and YY males in mixed populations (Franco *et al.*, 1982), implying that 25 years ago F^D (or a similar genetic element) was present across the entire range of Italy, as it is now. However, we did find F^D in Switzerland, where it was not detected before 1982, suggesting that the F^D factor has spread northwards slightly.

Table 5. *Logistic mixed-model analysis of the frequencies of XX males in the study of Franco et al. (1982) and this study*

Model	DF	Deviance	F	P
Population (random) + study	13	107.7		
Population (random)	14	117.5	1.19	0.7

The full model includes population as a random effect and study (data from Franco *et al.*, 1982 or from our study) as a fixed effect under analysis.

No significant difference between studies was found.

4. Discussion

Our results show that autosomal *M* factors have not spread northwards in Europe over the last 25 years, in contrast to what was predicted by Franco *et al.* (1982). One may argue that we have overlooked low frequencies of autosomal *M* factors in Switzerland and Germany due to insufficient sample size. Although this may be true, very low frequencies of autosomal factors still support the hypothesis that the standard XY system is not being replaced by autosomal factors in northern populations. In line with our results, we suggest that after their initial spread in southern localities (see Franco *et al.*, 1982), autosomal *M* factors reached a stable distribution.

Our results indicate that some factors prevent the spread of autosomal *M* in populations north of Italy. In the transect we studied, the Alps may be considered as a barrier, although the biology of the housefly and its ease of spread with human transportation seem to preclude this physical barrier as being important for the potential long-term spread of autosomal *M* factors. In fact, the presence of the F^D factor north of the Alps and the *M* factor on autosome II in flies collected in eastern France in 2004 (results not shown) suggests that geographical barriers do not prevent the northward spread of autosomal *M* factors. More likely, some climatic factors are responsible for the stability of the distribution of *M*. The most obvious climatic factor related to latitude is temperature, which has been shown to be a strong predictor of the frequencies of different sex-determining factors in the housefly worldwide (Feldmeyer *et al.*, submitted). However, it is not obvious how temperature might influence the evolution and distribution of SD mechanisms (discussed in detail in Feldmeyer *et al.*, submitted).

Our statistical analysis reveals an effect of temperature, but also a significant interaction between temperature and latitude on the frequency of autosomal SD factors (Table 4). The interaction stems from the fact that at higher latitudes temperature has a positive effect on the frequencies of autosomal SD factors, whereas the opposite pattern is present at

lower latitudes (not shown). This may suggest that autosomal SD factors reach the highest frequencies at intermediate temperatures. However, autosomal SD factors have been found at high frequencies in places where average temperatures are higher than at our sampling sites (Feldmeyer *et al.*, submitted). A more likely explanation is that temperature interacts with other climatic factors (such as humidity) that could be correlated with latitude (and altitude) in our study area. This could also explain why an *M* factor on autosome III and F^D have been found at locations in England where the yearly range of temperatures is similar to that in Germany and Switzerland (Denholm *et al.*, 1985; data on temperatures from WORLDCLIM, not shown). Additionally, *M* factors located on different autosomes may be differently affected by temperature.

It has also been proposed that autosomal *M* factors have spread due to their linkage with insecticide resistance genes (Kerr, 1970; Franco *et al.*, 1982), since the isolation of autosomal *M* factors coincided with the appearance of insecticide resistance in natural populations of the housefly (Tomita & Wada, 1989b). Also, in a number of resistant populations autosomal *M* males have been found (Tsukamoto, 1983) and one laboratory experiment showed replacement of standard XY males by autosomal *M* males after several generations of selection for DDT resistance (Kerr, 1970). However, even though linkage with insecticide resistance genes could facilitate spread of autosomal *M* factors, it is not clear how it could contribute to the clinal distribution of SD factors in the housefly. One could argue that in warmer climates more generations of flies are produced and more applications of insecticides are used, allowing faster spread of *M* factors linked with insecticide-resistant genes. However, since pesticides have been used throughout Europe for decades and resistance genes are widespread also in northern populations (Keiding, 1977, 1999), one would expect that *M* factors would be increasing in frequency, although more slowly, in the north also. As we showed in this study, this is not the case. Another argument is that there is no correlation between the frequency of autosomal *M* males and insecticide resistance in housefly populations from the eastern United States (Hamm *et al.*, 2005). Therefore, linkage with insecticide resistance genes might explain the spread of autosomal *M* factors in some cases, but it seems unlikely to provide a general explanation for the clinal distribution of SD factors in the housefly.

Interestingly, autosomal *M* factors are not fixed in most populations and multiple factors on several or even all chromosomes can be maintained in a single population. This polymorphism was one of the reasons underlying the opinion of earlier researchers that the sex-determining mechanism in the housefly is in a transient state (e.g. Franco *et al.*, 1982; Denholm

et al., 1985; Tomita & Wada, 1989b). However, theoretical models reveal that such a polymorphism can be stable not only for specific fitness values of different genotypes (Bull & Charnov, 1977; Jayakar, 1987), but also when different genotypes have the same viability and fertility (Kozielska *et al.*, 2006). Therefore, the conditions for a stable polymorphism may be much less restrictive than previously thought, and it may well be that the multifactorial SD system of the housefly is stable.

Unfortunately, we do not have data on the frequencies of different autosomal *M* factors in the past to see whether these frequencies have changed. Franco and colleagues (1982) did not find any *M* factors located on autosomes I, IV or V, but they do not provide the number of males investigated. If these factors were present in the past at low frequencies as they are now (Table 3), Franco *et al.* (1982) might not have detected them in small sample sizes. They reported that *M* was more common on autosome III than on autosome II. The same pattern is seen in this study and several others (Tomita & Wada, 1989b; Denholm *et al.*, 1990; Hamm *et al.*, 2005; except for Tanzanian populations, Feldmeyer *et al.*, submitted.). This suggests that *M* on autosome III confers the largest fitness gain to its bearer, but this may only be a conditional effect (e.g. frequency- or temperature-dependent) since the *M* on autosome III did not replace other *M* factors in recent decades in the Italian populations.

Another explanation for the high polymorphism in genomic location of *M* factors is that the *M* factor is part of a transposable element, as is known for the *M* factor in *Megaselia scalaris* (Traut & Willhoeft, 1990). In this species transposition rate differs depending on the chromosome on which *M* is located (Green, 1980). This might not only explain why *M* factors are more common on some autosomes than others, but also the clinal distribution of *M* factors, since transposition rate is known to be dependent on temperature and often increases with increasing temperature (Lampe *et al.*, 1998; Ohtsubo *et al.*, 2005; but see Hashida *et al.*, 2003). Molecular studies are necessary to establish whether the *M* factor is always the same gene located on a transposable element or whether *M* factors on different chromosomes are different genes blocking the female-determining factor *F* (see Dübendorfer *et al.*, 2002).

Our crosses suggest that the frequency of the Y chromosome has increased over recent decades in some Italian populations. We found an *M* factor on the sex chromosomes in some populations that were described as purely autosomal by Franco and colleagues (1982; Fig. 3). It is difficult to assess what the cause of these changes in particular populations is; some local factors may be involved. For population IT5, the difference between past and present

frequencies of XX males might reflect the fact that we could not locate accurately the sampling site of Franco and colleagues (1982; see Section 2). Moreover, it should be noted that due to the absence of visible markers on the sex chromosomes of the housefly, our crosses did not allow us to determine whether the *M* factor was present on the Y or on the X chromosome (as found in England: Denholm *et al.*, 1983, 1985). Without additional information, the data obtained from the crosses could easily lead to the incorrect classification of XX^{*M*} males as XY males. Therefore, we performed additional cytological investigations, using orcein staining, a standard technique employed in cytological studies of the housefly (Hiroyoshi, 1964; Franco *et al.*, 1982; Denholm *et al.*, 1983, 1985). Our preliminary results (not shown) confirm that males from the northernmost populations (GE1, GE2, SW and IT3) are of karyotype XY. Unfortunately, we could not unambiguously distinguish between XX, XY and YY karyotypes in the other populations, because the length polymorphism of the housefly sex chromosomes (also known from other strains: Boyes *et al.*, 1964; Boyes, 1967; Milani, 1971; Franco *et al.*, 1982; Denholm *et al.*, 1983, 1985; Hediger *et al.*, 1998) did not allow a reliable distinction between X and Y chromosomes. Therefore, we cannot exclude the possibility that the X chromosome (rather than the Y chromosome) bears the *M* factor in the southern populations.

In conclusion, even if the distribution of the Y chromosome in European populations is difficult to assess, our main conclusion that autosomal *M* factors have not spread northwards in the last 25 years still holds. This suggests that the polymorphism of SD factors in natural housefly populations is not transient but stable. Additional studies, at both the ecological and the molecular level, are required to unravel the factors responsible for the stable co-existence of various SD factors. Undoubtedly, better understanding of the housefly SD system will also provide general insights into the evolution of sex determination, which is still poorly understood in other taxa as well.

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